UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450

NOTICE OF ALLOWANCE AND FEE(S) DUE

7590

11/05/2003

JOSEPH LUCCI, ESQ. WOODCOCK WASHBURN LLP ONE LIBERTY PLACE 46TH FLOOR PHILADELPHIA, PA 19103

EXAMINER	
 FAV ZOHREH A	

ART UNIT PAPER NUMBER

1614

DATE MAILED: 11/05/2003

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/802,709	03/08/2001	Andrew C. Lam	ARC2865N1	1161

TITLE OF INVENTION: METHODS AND DEVICES FOR PROVIDING PROLONGED DRUG THERAPY

APPLN, TYPE	SMALL ENTITY	ISSUE FEE	PUBLICATION FEE	TOTAL FEE(S) DUE	DATE DUE
nonprovisional	NO	\$1330	\$300	\$1630	02/05/2004

THE APPLICATION IDENTIFIED ABOVE HAS BEEN EXAMINED AND IS ALLOWED FOR ISSUANCE AS A PATENT. PROSECUTION ON THE MERITS IS CLOSED. THIS NOTICE OF ALLOWANCE IS NOT A GRANT OF PATENT RIGHTS. THIS APPLICATION IS SUBJECT TO WITHDRAWAL FROM ISSUE AT THE INITIATIVE OF THE OFFICE OR UPON PETITION BY THE APPLICANT. SEE 37 CFR 1.313 AND MPEP 1308.

THE ISSUE FEE AND PUBLICATION FEE (IF REQUIRED) MUST BE PAID WITHIN THREE MONTHS FROM THE MAILING DATE OF THIS NOTICE OR THIS APPLICATION SHALL BE REGARDED AS ABANDONED. STATUTORY PERIOD CANNOT BE EXTENDED. SEE 35 U.S.C. 151. THE ISSUE FEE DUE INDICATED ABOVE REFLECTS A CREDIT FOR ANY PREVIOUSLY PAID ISSUE FEE APPLIED IN THIS APPLICATION. THE PTOL-85B (OR AN EQUIVALENT) MUST BE RETURNED WITHIN THIS PERIOD EVEN IF NO FEE IS DUE OR THE APPLICATION WILL BE REGARDED AS ABANDONED.

HOW TO REPLY TO THIS NOTICE:

I. Review the SMALL ENTITY status shown above.

If the SMALL ENTITY is shown as YES, verify your current SMALL ENTITY status:

- A. If the status is the same, pay the TOTAL FEE(S) DUE shown above.
- B. If the status is changed, pay the PUBLICATION FEE (if required) and twice the amount of the ISSUE FEE shown above and notify the United States Patent and Trademark Office of the change in status, or

If the SMALL ENTITY is shown as NO:

- A. Pay TOTAL FEE(S) DUE shown above, or
- B. If applicant claimed SMALL ENTITY status before, or is now claiming SMALL ENTITY status, check the box below and enclose the PUBLICATION FEE and 1/2 the ISSUE FEE shown above.
- ☐ Applicant claims SMALL ENTITY status. See 37 CFR 1.27.
- II. PART B FEE(S) TRANSMITTAL should be completed and returned to the United States Patent and Trademark Office (USPTO) with your ISSUE FEE and PUBLICATION FEE (if required). Even if the fee(s) have already been paid, Part B - Fee(s) Transmittal should be completed and returned. If you are charging the fee(s) to your deposit account, section "4b" of Part B - Fee(s) Transmittal should be completed and an extra copy of the form should be submitted.
- III. All communications regarding this application must give the application number. Please direct all communications prior to issuance to Mail Stop ISSUE FEE unless advised to the contrary.

IMPORTANT REMINDER: Utility patents issuing on applications filed on or after Dec. 12, 1980 may require payment of maintenance fees. It is patentee's responsibility to ensure timely payment of maintenance fees when due.

PART B - FEE(S) TRANSMITTAL

Complete and send this form, together with applicable fee(s), to: Mail

Mail Stop ISSUE FEE Commissioner for Patents P.O. Box 1450 Alexandria, Virginia 22313-1450

or Fax (703) 746-4000

INSTRUCTIONS: This form should be used for transmitting the ISSUE FEE and PUBLICATION FEE (if required). Blocks 1 through 4 should be completed where appropriate. All further correspondence including the Patent, advance orders and notification of maintenance fees will be mailed to the current correspondence address as indicated unless corrected below or directed otherwise in Block 1, by (a) specifying a new correspondence address; and/or (b) indicating a separate "FEE ADDRESS" for maintenance fee notifications.

CURRENT CORRESPONDENCE ADDRESS (Note: Legibly mark-up with any corrections or use Block 1)

7590

11/05/2003

JOSEPH LUCCI, ESQ. WOODCOCK WASHBURN LLP ONE LIBERTY PLACE 46TH FLOOR PHILADELPHIA, PA 19103

Note: A certificate of mailing can only be used for domestic mailings of the Fec(s) Transmittal. This certificate cannot be used for any other accompanying papers. Each additional paper, such as an assignment or formal drawing, must have its own certificate of mailing or transmission.

Certificate of Mailing or Transmission
I hereby certify that this Fec(s) Transmittal is being deposited with the United States Postal Service with sufficient postage for first class mail in an envelope addressed to the Mail Stop ISSUE FEE address above, or being facsimile transmitted to the USPTO, on the date indicated below.

	 _
(Depositor's name	
(Signature	
(Date	

APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/802,709	03/08/2001	Andrew C. Lam	ARC2865N1	1161

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nonprovisional	NO	\$1330		\$300	\$1630	02/05/2004
EXAM	MINER	ART UNI	Т	CLASS-SUBCLASS]	
FAY, ZO	OHREH A	1614		514-532000	_	•
Change of correspondence address or indication of "Fee Address" (37 CFR 1.363). Change of correspondence address (or Change of Correspondence Address form PTO/SB/122) attached.			names o agents O firm (hav	inting on the patent front page f up to 3 registered patent a R, alternatively, (2) the name ring as a member a registered the names of up to 2 regis	of a single attorney or 2	
☐ "Fee Address" indication (or "Fee Address" Indication form PTO/SB/47; Rev 03-02 or more recent) attached. Use of a Customer Number is required.				or agents. If no name is liste		

3. ASSIGNEE NAME AND RESIDENCE DATA TO BE PRINTED ON THE PATENT (print or type)

PLEASE NOTE: Unless an assignce is identified below, no assignce data will appear on the patent. Inclusion of assignce data is only appropriate when an assignment has been previously submitted to the USPTO or is being submitted under separate cover. Completion of this form is NOT a substitute for filing an assignment. (B) RESIDENCE: (CITY and STATE OR COUNTRY)

(A) NAME OF ASSIGNEE

Please check the appropriate assignee category	or categories (will not be printed on the patent);	☐ individual	□ corporation or ot	her private group entity	government
4a. The following fee(s) are enclosed:	4b. Payment of Fee(s):				
☐ Issue Fee	☐ A check in the an	nount of the fee(s)	is enclosed.		
□ Publication Fec	Payment by credi	it card. Form PTO-	2038 is attached.		
Advance Order - # of Copies	☐ The Director is Deposit Account No	hereby authorized umber	by charge the requir	ed fee(s), or credit any ose an extra copy of this	overpayment, to form).
Director for Patents is requested to apply the Is	ssue Fee and Publication Fee (if any) or to re-appl	y any previously p	oaid issue fee to the ap	plication identified abox	ye.
(Authorized Signature)	(Date)		w.		
NOTE; The Issue Fee and Publication Fee other than the applicant; a registered attorn interest as shown by the records of the United	(if required) will not be accepted from anyone ney or agent; or the assignee or other party in a States Patent and Trademark Office.				
estimated to take 12 minutes to complete, in completed application form to the USPTO. case. Any comments on the amount of ti suggestions for reducing this burden, should	y 37 CFR 1.311. The information is required to ch is to file (and by the USPTO to process) an 5 U.S.C. 122 and 37 CFR 1.14. This collection is cluding gathering, preparing, and submitting the Time will vary depending upon the individual me you require to complete this form and/or it be sent to the Chief Information Officer, U.S. partment of Commerce, Alexandria, Virginia COMPLETED FORMS TO THIS ADDRESS. Indria, Virginia 22313-1450.				
Under the Paperwork Reduction Act of 19 collection of information unless it displays a	995, no persons are required to respond to a valid OMB control number.				



United States Patent and Trademark Office

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450

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09/802,709	03/08/2001	Andrew C. Lam	ARC2865N1	1161
7	590 11/05/2003		EXAM	INER
JOSEPH LUCCI			FAY, ZO	HREH A
WOODCOCK WA			ART UNIT	PAPER NUMBER
46TH FLOOR			1614	· · · · ·
PHILADELPHIA, PA 19103			DATE MAILED: 11/05/200	3

Determination of Patent Term Adjustment under 35 U.S.C. 154 (b)

(application filed on or after May 29, 2000)

The Patent Term Adjustment to date is 0 day(s). If the issue fee is paid on the date that is three months after the mailing date of this notice and the patent issues on the Tuesday before the date that is 28 weeks (six and a half months) after the mailing date of this notice, the Patent Term Adjustment will be 0 day(s).

If a Continued Prosecution Application (CPA) was filed in the above-identified application, the filing date that determines Patent Term Adjustment is the filing date of the most recent CPA.

Applicant will be able to obtain more detailed information by accessing the Patent Application Information Retrieval (PAIR) system (http://pair.uspto.gov).

Any questions regarding the Patent Term Extension or Adjustment determination should be directed to the Office of Patent Legal Administration at (703) 305-1383. Questions relating to issue and publication fee payments should be directed to the Customer Service Center of the Office of Patent Publication at (703) 305-8283.



UNITED STATES PATENT AND TRADEMARK OFFICE

UNITED STATES DEPARTMENT OF COMMERCE United States Patent and Trademark Office Address: COMMISSIONER FOR PATENTS P.O. Box 1450 Alexandria, Virginia 22313-1450

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09/802,709	03/08/2001	Andrew C. Lam	ARC2865N1	1161
7:	590 11/05/2003		EXAM	INER
JOSEPH LUCCI			FAY, ZO	HREH A
WOODCOCK WA			ART UNIT	PAPER NUMBER
46TH FLOOR	LACE		1614	
PHILADEL PHIA	PA 19103			

Notice of Fee Increase on October 1, 2003

If a reply to a "Notice of Allowance and Fee(s) Due" is filed in the Office on or after October 1, 2003, then the amount due will be higher than that set forth in the "Notice of Allowance and Fee(s) Due" since there will be an increase in fees effective on October 1, 2003. See Revision of Patent Fees for Fiscal Year 2004; Final Rule, 68 Fed. Reg. 41532, 41533, 41534 (July 14, 2003).

The current fee schedule is accessible from (http://www.uspto.gov/main/howtofees.htm).

If the fee paid is the amount shown on the "Notice of Allowance and Fee(s) Due" but not the correct amount in view of the fee increase, a "Notice of Pay Balance of Issue Fee" will be mailed to applicant. In order to avoid processing delays associated with mailing of a "Notice of Pay Balance of Issue Fee," if the response to the Notice of Allowance is to be filed on or after October 1, 2003 (or mailed with a certificate of mailing on or after October 1, 2003), the issue fee paid should be the fee that is required at the time the fee is paid. If the issue fee was previously paid, and the response to the "Notice of Allowance and Fee(s) Due" includes a request to apply a previously-paid issue fee to the issue fee now due, then the difference between the issue fee amount at the time the response is filed and the previously-paid issue fee should be paid. See Manual of Patent Examining Procedure, Section 1308.01 (Eighth Edition, August 2001).

Effective October 1, 2003, 37 CFR 1.18 is amended by revising paragraphs (a) through (c) to read as set forth below.

Section 1.18 Patent post allowance (including issue) fees.

(a) Issue fee for issuing each original or reissue patent, except a design or plant patent:

(b) Issue fee for issuing a design patent:

(c) Issue fee for issuing a plant patent:

By a small entity (Sec. 1.27(a))......\$320.00

By other than a small entity......\$640.00

Questions relating to issue and publication fee payments should be directed to the Customer Service Center of the Office of Patent Publication at (703) 305-8283.

-		
	Application No.	Applicant(s)
Netice of Allowshiller	09/802,709	LAM ET AL.
Notice of Allowability	Examiner	Art Unit
	Zohreh Fay	1614
<u> </u>		
The MAILING DATE of this communication apperature All claims being allowable, PROSECUTION ON THE MERITS IS therewith (or previously mailed), a Notice of Allowance (PTOL-85) NOTICE OF ALLOWABILITY IS NOT A GRANT OF PATENT RIPORT of the Office or upon petition by the applicant. See 37 CFR 1.313	(OR REMAINS) CLOSED in this app or other appropriate communication GHTS. This application is subject to	olication. If not included will be mailed in due course. THIS
1. This communication is responsive to the amendments and	remarks filed on Octoer 15, 2003.	
2. The allowed claim(s) is/are <u>37 and 46-50</u> .		
3. \boxtimes The drawings filed on <u>08 March 2001</u> are accepted by the	Examiner.	
4. Acknowledgment is made of a claim for foreign priority und	er 35 U.S.C. § 119(a)-(d) or (f).	
a) All b) Some* c) None of the:		•
 Certified copies of the priority documents have 	been received.	
Certified copies of the priority documents have	been received in Application No	·
Copies of the certified copies of the priority doc	uments have been received in this r	national stage application from the
International Bureau (PCT Rule 17.2(a)).		
* Certified copies not received:		
5. Acknowledgment is made of a claim for domestic priority un	• , , ,	onal application).
(a) The translation of the foreign language provisional ap	, •	$p^* \phi = p^*$
6. Acknowledgment is made of a claim for domestic priority un	der 35 U.S.C. §§ 120 and/or 121.	
Applicant has THREE MONTHS FROM THE "MAILING DATE" of below. Failure to timely comply will result in ABANDONMENT of to 7. A SUBSTITUTE OATH OR DECLARATION must be subminformal patent Application (PTO-152) which gives reason	his application. THIS THREE-MON itted. Note the attached EXAMINER	ITH PERIOD IS NOT EXTENDABLE. 'S AMENDMENT or NOTICE OF
· · · · · · · · · · · · · · · · · · ·	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	
 8. CORRECTED DRAWINGS must be submitted. (a) including changes required by the Notice of Draftspers 1) hereto or 2) to Paper No. 	on's Patent Drawing Review (PTO-	948) attached
(b) \square including changes required by the proposed drawing c	orrection filed $_{}$, which has be	een approved by the Examiner.
(c) including changes required by the attached Examiner's	Amendment / Comment or in the C	Office action of Paper No
Identifying indicia such as the application number (see 37 CFR 1.6 each sheet.	B4(c)) should be written on the drawin	ags in the front (not the back) of
 DEPOSIT OF and/or INFORMATION about the depose attached Examiner's comment regarding REQUIREMENT FOR THE CONTROL OF THE PROPERTY OF THE PROPERTY	it of BIOLOGICAL MATERIAL m HE DEPOSIT OF BIOLOGICAL MAT	nust be submitted. Note the FERIAL.
Attachment(s)		
 1 Notice of References Cited (PTO-892) 3 Notice of Draftperson's Patent Drawing Review (PTO-948) 5 Information Disclosure Statements (PTO-1449), Paper No	4☐ Interview Summa 6☐ Examiner's Amer	ary (PTO-413), Paper No Indment/Comment Iment of Reasons for Allowance Indiany Examiner CROUP 1200

	ALLOWED REVIEW		ε ² - γ	*
Application Number 09/802,709	Notice Of Allowance 05-Nov-03	Examiner FAY, ZOHREH A		
1 · · · · ·	SECTION XI. File Wrapper			
Reasons for Allowance (R/	/A)	,		
•	Reasons for Allowance (R/A)?	\circ Yes	No No	$\circ_{N/A}$
If yes,				
Is the R/A clear and co	mplete?	\circ Yes	\circ_{No}	
If no,				
Does the record as a w	hole indicate a R/A was necessary?	O Yes	No	
Comments:				
Interviews				
Was there an interview regaction reviewed?	garding the merits of the case relevant to the	○ _{Yes}	⊙ _{No}	
If yes,	·			
Was Summary Form PTOL-	•	O Yes		
Is the record of the inte	erview clear and complete?	○ Yes	○ No	
Claims		-	_	
•	nappropriate manner on non-substantive issues?	O Yes	No	
If yes, \Box claims are present tha	st ware not addressed			
		-111		
<u> </u>	drawn from consideration should have been cance	illea.		
	laims were not properly treated.			
\square other				
Comments:				
Sequence Rules				
Does the application contain	nucleotide and/or amino acid sequences?	\circ Yes	No No	
If yes,		_	_	
Did the examiner properly h	andle Sequence Compliance Issues?	○ Yes	O No	
Comments:				

	ALLOWED REVIEW		
Application Number 09/802,709	Notice Of Allowance 05-Nov-03	Examiner FAY, ZOHREH A	
Se	ction XIV. Indicia of Commendable	e/Outstanding	1
Patentability Determination	n: Indicia of Commendable/Outst	anding	
	the examiner shows an indication of a sistent with the file record and prosec	•	□Yes
	d arguments made by the examiner, d which results in amendment(s) prop		□Yes
	application clearly shows that the exa dest reasonable interpretation and se elated art areas.		□Yes
Action Taking: Indicia of Co	mmendable/Outstanding		
	objection, and response to argument nded in the resulting Office actions ind positions to the applicant.		□Yes
•	er an applicant's attention to relevant Office action relies to support the pos		□Yes
the principle of compact prospossible including consultation (see search guidelines); plactilisms as well as other art with disclosed invention; and issu	nat the principle of compact prosecution comprises conducting an inition with an expert in the art where the ing art of record which meets both the hich is pertinent to significant thoughing a first Office action which clearly children detail that absent some unexpected.	ial search which is as complete as e examiner lacks such expertise ne concept and the wording of the unclaimed features of the explains the examiner's position	□Yes
Patent Examining Function:	Indicia of Commendable/Outsta	inding	
complete file wrapper r	tatement if applicable: mulated to advance the prosecution, ecord. The Office action also is such ice action clearly and concisely preser	that it leaves little room for	lop a
	is complete and accurate and does reconveys the positions taken.	not require any substantial revision.	The
Comments:			
	·		
	-		

PLEASE DELIVER TO:

GWEN PAYNE CM1-7D11

ALLOWED REVIEW POTENTIAL CLEAR ERROR

REVIEWER - ATTACH TO APPLICATION

		3.	ALLOWED RE	EVIEW	3		
Application Number 09/802,709	Art Unit 1614	Notice of Allowance 05-Nov-03	2			Examin FAY, ZOHR	
4 4		·	Omitted Reje	ctions	*,		Marie Arthritis (Arthri
Is there a pote you propose m			nitting a rejec	ction? (The	e rejection	•	Yes 🔾 No
If yes, check all							
☐ 35 U.S.C. 1	.03						
☑ 35 U.S.C. 1	.12, first para	graph, writte	en description				
□ 35 U.S.C. 1					•		
☐ 35 U.S.C. 1	112, second p	aragraph					
□ 35 U.S.C. 1							:
☐ 35 U.S.C. 1	.01 (non-stat	utory subjec	t matter)	•			
☐ Double Pat	enting (statut	ory, ODP)					
\square Other (e.g.	, Best Mode)						
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]

ALLOWED REVIEW Notice Of Allowance Application Number Examiner 09/802,709 05-Nov-03 FAY, ZOHREH A Search **Initial Data Capture Points** O Yes O No Was art provided from an ESS before first action? \bigcirc N/A Was any IDS improperly treated? ○ Yes • No Yes ○ No Was a text search performed by the Examiner? · • Yes O No Was it non-patent literature? \bigcirc N/A Yes ○ No Is the search strategy printout present? Was the inventorship searched by the Examiner? If any TC or specialized art has identified specific search requirements in ○ Yes ○ No N/A addition to or in exclusion of the above requirements, were these specific requirements complied with? ○ Yes • No Were foreign patent documents cited by the Examiner on an 892? O Yes No Were NPL documents cited by the Examiner on an 892? Yes ○ No. \bigcirc N/A Did the Examiner perform a new search in a 2nd/subsequent action? Yes ○ No \bigcirc N/A Did the Examiner update all searches in a subsequent action? ○ Yes ○ No \bigcirc N/A Did an ESS submit a new search report in a 2nd/subsequent action? Was there new art found by the Examiner that was applied in a O N/A Yes ○ No ■ 2nd/subsequent action? ○ Yes • No Has a search been performed by the Reviewer? **Overall Rating of the Search** • Adequate ○ Less than Adequate Comments:

	ALLOWED REVIEW	7		· · · · · · · · · · · · · · · · · · ·			
Application Number	Notice Of Allowance		Exam	niner			
09/802,709	05-Nov-03		FAY, ZOI	HREH A			
*	Section III. 35 U.S.C.	102					
Correctness of 35 U.S.C 10)2 Rejections						
Were all 35 U.S.C 102 reje	-		0 0	`			
(No indicates potenti	○ Yes	⁾ No					
If no, indicate the problem							
☐ Claimed features no	t found in the reference.						
☐ Wrong subsection of	35 U.S.C. 102 used.						
☐ Date of the reference	e no good.						
\square Inherency applied in	_						
☐ Improper official not							
☐ Other							
Comments:				,			
Clarity of 35 U.S.C 102 Rej	ections	0					
	ections ections formulated in a clear manne	~~?	○ Yes ○	. Ni-			
(No indicates potentia		31 f	○ res ○	INO			
Were claim limitations mat				_ ,			
		_	Sometimes		\circ		
Was any statement of inhe Comments:	rency clearly explained?	\cdot Yes	O Sometimes	○ No	O _{N/A}		
Comments.	· .						
·							
	That Should Have Been Made						
Give a brief description of t	he proposed35 U.S.C. 102 rejection	ns(s) that sl	hould have been	made:			
	e rejected under 35 USC 102(b) as	•	over J. W. Hubb	oard et a	al (J. of		
	s, Vol 78, No. 11, November 1989,		J (ADD) ALL	L'	.		
The claims are drawn to a method for treating Attention-Deficit Disorder (ADD) or Attention-Deficit							
	Hyperactivity Disorder (ADHD) in a patient comprising administering a pharmaceutically acceptable composition comprising 100 ng to 500 mg of methylphenidate (MPH) to achieve a-substantially						
ascending methylphenidate plasma drug concentration over a time period of about 5.5 hours (claim							
37). The specification (page 16, lines 1-8) defines "ascending release rate" as "a periodic release rate							
that is increased over the immediately-preceding periodic release rate, where the periodic intervals							
are the same. For example, when the quantity of drug released from a dosage form is measured at							
hourly intervals and the quantity of drug released during the fifth hour following administration (determined at t=5hours) is greater than the quantity of drug released from the dosage form during							
the fourth hour following administration (determined at t=4 hours), an ascending release rate from							
the fourth hour to the fifth hour has occurred."							
Hubbard et al teach the administration of methylphenidate is widely used to treat children with ADD							
(page 944) and that in this study all children were administered 20 mg of MPH-SR (sustained							
release). The curve representing I-MPH from Patient profile 2 teaches an ascending sustained plasma concentration up to 6 hours. The claims contain the language "substantially ascending							
methylphenidate plasma drug concentration", while the specification technically only defines							
"ascending release rate"	The specification (page 4, lines 6	-13) disclo	ses "It is believe	d to be			
particularly desirable to	provide sustained release oral dosa	age forms tl	hat provide drug	release			
substantially constant release rate over an extended time period. In this manner, for many drugs, the							

ALLOWED REVIEW Notice Of Allowance **Application Number Examiner** 09/802,709 05-Nov-03 FAY, ZOHREH A Section V. 35 U.S.C. 112 1st paragraph, written description Correctness of 35 U.S.C 112 1st Paragraph, Written Description Rejections Were all 35 U.S.C 112 1st paragraph, written description rejections Oyes O No reasonable? (No indicates potential clear error) Comments: Clarity of 35 U.S.C 112 1st Paragraph, Written Description Rejections Were all 35 U.S.C 112 1st paragraph, written description rejections Oyes O No formulated in a clear manner? Comments: 35 U.S.C 112 1st Paragraph, Written Description Rejection(s) That Should Have Been Made Give a brief description of the 112 1st paragraph, written description rejections that should have been Claims 37, 49 and 50 (renumbered under Rule 126) are rejected under 35 USC 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention. Vas-Cath Inc v. Mahurkar, 19 USPQ2d 1111 (Fed. Cir. 1991), clearly states that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the 'written description' inquiry, whatever is now claimed." Vas-Cath Inc. v. Mahurkar, 19USPQ2d at 1117. Applicant's invention is drawn to a method for treating ADD or ADHD in a patient, wherein the method comprises administering a pharmaceutically acceptable composition comprising 100 ng to 500 mg of methylphenidate and a pharmaceutically acceptable carrier to said patient in a manner that achieves a substantially ascending methylphenidate plasma drug concentration over a time period of about 5.5 hours following said administration. However, the claimed concentration of 100 ng to 500 mg is not disclosed within the instant specification. These claims were submitted by preliminary amendment but the declaration fails to refer to the submission of any preliminary amendment. The originally filed claims do not include the above noted concentration. Applicant might consider petitioning this issue and submitting a new declaration referring to the preliminary amendment in order to resolve the written description problem. In the meantime, the specific concentration of 100 ng to 500 mg of methylphenidate in claims 37, 49 and 50 is found to have no support in the instant specficiation.

A. (ALLOWED REVIEW		1 ×	
Application Number	Notice Of Allowance		Exami	ner
09/802,709 05-Nov-03		FAY, ZOHREH A		
	Section XIII. Other Issues	12 ¹ 0 4		
Were all claims for priority properly treated?			\circ_{No}	$\circ_{N/A}$
If a restriction was made, was it proper?			\circ_{No}	N/A
Were all matters of substance in applicant's response and affidavits/declarations evaluated sufficiently?			No	○ _{N/A}
Other issues?		\circ_{Yes}	⊙ No	
Applicant's argument regard examiner.	ling the Hubbard et al reference was not c	orrectly eval	luated by	the

Comments:

If the rejection under 35 USC 102 is agreed upon, the Examiner should additionally set forth the following ODP rejection. Claims 37 and 46-50 provisionally rejected under the judicially created doctrine of obviousness-type double patenting as being unpatentable over claims 15, 88 and 90-95 of copending application no. 09/253,317. Although the conflicting claims are not identical, they are not patenably distinct from each other because they are both claiming a method for treating ADD or ADHD comprising administering methylphenidate at an ascending release rate. The dosage form comprises the same range of 100 ng to 500 mg and the time periods are equally covered. This is a provisional obviousness-type double patenting rejection because the conflicting claims have not in fact been patented. The instant application 09/802,709 is a continuation of 09/253,317.

plasma drug concentration initially ascends for a short period of time as drug release begins and then remains substantially constant over an extended time period as drug release continues at a constant rate." Looking at the period of time for Patient Profile 2 between 6 and 12 hours the I-MPH is declining, but the level still remains consistently higher than the plasma level prior to the initial administration of MPH-SR. Although not included in this rejection, claims 49 and 50 could be considered as anticipated by Hubbard et al, in view of the case law to follow for interpreting "substantially". Hubbard et al teach "The plot of mean data (Figure 2) further supports this observation of sustained plasma concentration for both isomers up to 6 hours." (page 945) Hubbard concludes by stating "this pilot study demonstrated that despite higher plasma levels of d-MPH, the levels of both the enantiomers were sustained for over a period of 8 hours in all the six children with ADD or ADD-H dosed with 20 mg of MPH-SR." (page 947) In view of the inclusion of "sustained for over a period of 8 hours", further consideration should be given to claims 49 and 50 for this rejection.

A recent CAFC decision addresses the word "substantially" as not making a claim indefinite merely because it scope is not ascertainable from the face of the claims. (Amgen Inc. v. Hoechst Marion Roussel Inc., 65 USPQ2s 1385 (CAFC 2003), 1406) Thus, to properly understand and interpret "substantially" one should look to the specification for an understanding. In this case, page 4, lines 6-13 (as discussed above) of the instant specification provide the understanding that the drug is to be released at a substantially constant rate over an extended time period, thus providing an understanding for "substantially ascending methylphenidate plasma drug concentration over a time period".

The Examiner applied the Hubbard et al reference against claims 37 and 46-50 in paper #21 but referenced Patient Profile #3 which at best demonstrates an ascending release rate of 3-4 hours, if "substantially ascending" can be interpreted as still a higher plasma level than the initial administration level. The attorney argued that Profile #3 fails to support "ascending plasma drug formulation up to about 10 hours", which appears to be a correct assertion. Profile #3 does not meet the claim limitations, but Profile #2 does anticipate the instantly claimed invention for at least claims 37 and 46-48.

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Comments:			

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